

## Freeform Search

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<b>Database:</b>	<input checked="" type="checkbox"/> US Pre-Grant Publication Full-Text Database <input checked="" type="checkbox"/> US Patents Full-Text Database <input checked="" type="checkbox"/> <b>US OCR Full-Text Database</b> <input type="checkbox"/> EPO Abstracts Database <input type="checkbox"/> JPO Abstracts Database <input type="checkbox"/> Derwent World Patents Index <input type="checkbox"/> IBM Technical Disclosure Bulletins
<b>Term:</b>	<input type="text" value="L23 not 122"/> <span style="float: right; margin-top: -20px;"> <input checked="" type="checkbox"/>  <input type="checkbox"/> </span>
<b>Display:</b>	<input type="text" value="200"/> Documents in <u>Display Format:</u> [-] Starting with Number <input type="text" value="1"/>
<b>Generate:</b> <input type="radio"/> Hit List <input checked="" type="radio"/> Hit Count <input type="radio"/> Side by Side <input type="radio"/> Image	

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### Search History

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DATE: Thursday, October 05, 2006    [Purge Queries](#)    [Printable Copy](#)    [Create Case](#)

<u>Set</u>	<u>Hit</u>	<u>Set</u>
<u>Name</u>	<u>Count</u>	<u>Name</u>
side by side	result set	
<i>DB=PGPB,USPT,EPAB,JPAB,DWPI; PLUR=YES; OP=OR</i>		
<u>L24</u> L23 not l22	65	<u>L24</u>
<u>L23</u> sucralose same tablet and dextrose	87	<u>L23</u>
<u>L22</u> L21 and dextrose	22	<u>L22</u>
<u>L21</u> sucralose same tablet same (soft or chewable or chewing or disintegrat\$4)	41	<u>L21</u>
<u>L20</u> sucralose same tablet	140	<u>L20</u>
<u>L19</u> L18 and dextrose	17	<u>L19</u>
<u>L18</u> (fat or oil or lipid or triglyceride or glyceride) adj2 free same tablet	54	<u>L18</u>
<u>L17</u> (fat or oil or lipid or triglyceride or glyceride) adj3 free adj5 tablet	2	<u>L17</u>
<u>L16</u> (fat or oil or lipid or triglyceride or glyceride) adj3 free same tablet	61	<u>L16</u>
<u>L15</u> (fat or oil or lipid or triglyceride or glyceride) adj3 free same soft adj2 tablet	0	<u>L15</u>
<u>L14</u> chewable adj2 tablet same (sugar or non-sweet\$4 or sweet) adj3 free	18	<u>L14</u>
<u>L13</u> (fat or oil or lipid or triglyceride or glyceride) adj3 free same chewable adj2 tablet	3	<u>L13</u>
<u>L12</u> (fat or oil or lipid or triglyceride or glyceride) adj3 free and chewable adj2 tablet same (sugar or non-sweet\$4 or sweet) adj3 free	2	<u>L12</u>

<u>L11</u>	(fat or oil or lipid or triglyceride or glyceride) adj3 free same chewable adj2 tablet and (sugar or non-sweet\$4 or sweet) adj3 free	2	<u>L11</u>
<u>L10</u>	(fat or oil or lipid or triglyceride or glyceride) adj3 free same tablet and (sugar or non-sweet\$4 or sweet) adj3 free same tablet	6	<u>L10</u>
<u>L9</u>	(fat or oil or lipid or triglyceride or glyceride) adj3 free and (sugar or non-sweet\$4 or sweet) adj3 free same tablet	9	<u>L9</u>
<u>L8</u>	(fat or oil or lipid or triglyceride or glyceride) adj3 free adj5 (sugar or non-sweet\$4 or sweet) same tablet	0	<u>L8</u>

*DB=PGPB,USPT; PLUR=YES; OP=OR*

<u>L7</u>	6596311.pn.	1	<u>L7</u>
<u>L6</u>	luber-joseph.in.	11	<u>L6</u>
<u>L5</u>	luber-j.in.	0	<u>L5</u>
<u>L4</u>	bunick-frank.in.	1	<u>L4</u>
<u>L3</u>	L2	45	<u>L3</u>

*DB=PGPB,USPT,EPAB,JPAB,DWPI; PLUR=YES; OP=OR*

<u>L2</u>	bunick.in.	131	<u>L2</u>
<u>L1</u>	bunick-f.in.	1	<u>L1</u>

END OF SEARCH HISTORY

=> d 11 ibib kwic

L1 ANSWER 1 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 2005:1193308 CAPLUS  
 DOCUMENT NUMBER: 143:466159  
 TITLE: Controlled release mucoadhesive matrix formulation containing tolterodine and a process for its preparation  
 INVENTOR(S): Durga Maheswari, Parvataneni; Appalaswamy Naidu, Rongala; Podile, Khadgapathi; Venkaiah Chowdary, Nannapaneni  
 PATENT ASSIGNEE(S): Natco Pharma Limited, India  
 SOURCE: PCT Int. Appl., 36 pp.  
 CODEN: PIXXD2  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE -
WO 2005105036	A1	20051110	WO 2005-IN99	20050404
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				

PRIORITY APPLN. INFO.: IN 2004-CH393 A 20040428  
 REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

AB Controlled release oral pharmaceutical mucoadhesive matrix formulation containing a therapeutically effective amount of tolterodine or its pharmaceutically acceptable salts, prodrugs and metabolites thereof dispersed in a rate controlling polymeric matrix comprising (1) a pH independent gelling polymer, such as polyethylene oxide, (2) pH dependent gelling polymer, such as sodium CM-cellulose (3) a film coating polymer component, such as Eudragit RS100 and other conventional tablet functional excipients. The formulation such as tablets or minitablets in capsules of the present invention relates to a 24 h controlled release dosage form useful for the treatment of urge incontinence and other symptoms of unstable or overactive urinary bladder. The invention also relates to a process for the preparation of controlled release mucoadhesive matrix formulation containing tolterodine in a tablet or mini tablets in capsule dosage form. For example, controlled-release mucoadhesive matrix tablets were prepared by wet granulation of tolterodine tartrate 2.0, polyethylene oxide-18 NF 7.0, sodium CM-cellulose 3.0, lactose anhydrous 20.0, microcryst. cellulose 51.8, polyvinylpyrrolidone K-30 5.0, Eudragit RS 100 10.0, iso-Pr alc. 72, and acetone 48, granules obtained were dried, lubricated

with colloidal silica 0.1, talc 0.1, and magnesium stearate 1.0 mg, resp., and compressed into core tablets. Eudragit L 100-55 3.0 was added to a mixture of iso-Pr alc. 25.6 and acetone 37.8, followed by tri-Et citrate 0.5 mg, resp., and the solution obtained was used as a barrier coating for core tablets.

IT Polysaccharides, biological studies  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(acidic; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Drug delivery systems  
(bioadhesive, mucoadhesive; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Drug delivery systems  
(capsules, controlled-release, minitablets-containing; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Alcohols, biological studies  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(fatty; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Cereal (grain)  
(hydrolyzed, solids; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Bladder, disease  
(incontinence, treatment of; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Polyesters, biological studies  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(lactic acid-based; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Polymers, biological studies  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(matrix; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Drug delivery systems  
(oral, controlled-release; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Bladder, disease  
(overactive bladder, treatment of; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Coating materials  
(polymer film; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Dissolution  
Gelation agents  
Gums and Mucilages  
Plasticizers  
(preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Adhesins  
Agglutinins and Lectins  
Bentonite, biological studies

Biopolymers  
Carbohydrates, biological studies  
Clays, biological studies  
Gelatins, biological studies  
Glycoproteins  
Hydrocarbon oils  
Kaolin, biological studies  
Polyesters, biological studies  
Polyoxyalkylenes, biological studies  
Smectite-group minerals  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Drug delivery systems  
(tablets, controlled-release; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Fats and Glyceridic oils, biological studies  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(vegetable, hydrogenated; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Granulation  
(wet; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 9003-01-4D, crosslinked  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(Carbopol; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 9003-39-8D, crosslinked  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(Crospovidone; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 9010-88-2, Eudragit NE 30D  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(Eudragit NE 50D; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 9050-36-6, Maltodextrin  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(Mor-Rex; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 7631-86-9, Silica, biological studies  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(colloidal; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 9004-34-6, Cellulose, biological studies  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(microcryst.; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 124937-51-5, Tolterodine 124937-52-6, Tolterodine tartrate  
RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 50-70-4, Sorbitol, biological studies 50-99-7, D-Glucose, biological studies 56-40-6, Glycine, biological studies 57-11-4, Stearic acid,

biological studies 57-48-7, Fructose, biological studies 57-50-1, Sucrose, biological studies 69-65-8, D-Mannitol 77-89-4, Acetyl triethyl citrate 77-90-7, Acetyl tributyl citrate 79-41-4D, Methacrylic acid, derivs., polymers 84-66-2, Diethyl phthalate 84-74-2, Dibutyl phthalate 87-89-8, Inositol 88-99-3, Phthalic acid, biological studies 102-76-1, Triacetin 108-32-7, Propylene carbonate 109-43-3, Dibutyl sebacate 112-92-5, Stearyl alcohol 117-81-7, Diethyl phthalate 134-03-2, Sodium ascorbate 471-34-1, Calcium carbonate, biological studies 557-04-0, Magnesium stearate 557-05-1, Zinc stearate 585-86-4, Lactitol 1327-43-1, Magnesium aluminum silicate 1344-95-2, Calcium silicate 1592-23-0, Calcium stearate 4070-80-8, Sodium stearyl fumarate 7789-77-7, Dibasic calcium phosphate dihydrate 9000-01-5, Acacia gum 9000-07-1, Carrageenan 9000-11-7, Carboxymethyl cellulose 9000-30-0, Guar gum 9000-36-6, Karaya gum 9000-40-2, Locust bean gum 9000-65-1, Tragacanth gum 9000-69-5, Pectin 9002-18-0, Agar 9002-88-4D, Polyethylene, alkyl ethers 9003-01-4, Poly(acrylic acid) 9003-39-8, Polyvinylpyrrolidone 9004-32-4, Sodium CM-cellulose 9004-38-0, CAP 9004-53-9, Dextrin 9004-57-3, Ethyl cellulose 9004-61-9, Hyaluronic acid 9004-62-0, Hydroxyethyl cellulose 9004-64-2, Hydroxypropyl cellulose 9004-65-3, Hypromellose 9004-67-5, Methyl cellulose 9005-25-8, Starch, biological studies 9005-32-7, Alginic acid 9005-38-3, Sodium alginate 9005-65-6, Polysorbate 80 9005-82-7, Amylose 9012-76-4, Chitosan 9050-31-1, HPMCP 9063-38-1, Sodium starch glycolate 10101-41-4, Calcium sulfate dihydrate 12705-30-5, Celutab 13463-67-7, Titanium dioxide, biological studies 14807-96-6, Talc, biological studies 18662-40-3, Calcium sulfate monohydrate 25086-15-1, Eudragit S 100 25212-88-8, Eudragit L 100-55 25322-68-3, Polyethylene oxide 25496-72-4, Glyceryl monooleate 26009-03-0, Poly(glycolic acid) 26023-30-3, Poly[oxy(1-methyl-2-oxo-1,2-ethanediyl)] 26100-51-6, Poly(DL-lactic acid) 26124-68-5, Poly(glycolic acid) 26936-24-3, Eudragit FS 30D 31566-31-1, Glyceryl monostearate 33434-24-1, Eudragit RS 100 36653-82-4, Cetyl alcohol 39301-46-7, Calcium pectinate 53237-50-6 66828-18-0, Dextrose 71138-97-1, HPMCAS 74811-65-7, Croscarmellose sodium 77538-19-3, Glyceryl behenate 77938-63-7, Dextrose monohydrate 139061-06-6, Calcium lactate trihydrate 147335-38-4, Eudragit NE 40D 178806-61-6, Eudragit RLPO 476312-12-6, Carbopol 71G 869094-48-4, Maltrons  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

=> d 11 ibib kwic 1-

YOU HAVE REQUESTED DATA FROM 23 ANSWERS - CONTINUE? Y/(N):y

L1 ANSWER 1 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 2005:1193308 CAPLUS  
 DOCUMENT NUMBER: 143:466159  
 TITLE: Controlled release mucoadhesive matrix formulation containing tolterodine and a process for its preparation  
 INVENTOR(S): Durga Maheswari, Parvataneni; Appalaswamy Naidu, Rongala; Podile, Khadgapathi; Venkaiah Chowdary, Nannapaneni  
 PATENT ASSIGNEE(S): Natco Pharma Limited, India  
 SOURCE: PCT Int. Appl., 36 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

1

## PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2005105036	A1	20051110	WO 2005-IN99	20050404
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				

PRIORITY APPLN. INFO.:

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RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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IT Polysaccharides, biological studies

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(acidic; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Drug delivery systems

(bioadhesive, mucoadhesive; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Drug delivery systems

- (capsules, controlled-release, minitablets-containing; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)
- IT Alcohols, biological studies  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(fatty; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)
- IT Cereal (grain)  
(hydrolyzed, solids; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)
- IT Bladder, disease  
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- IT Polymers, biological studies  
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(matrix; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)
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(oral, controlled-release; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)
- IT Bladder, disease  
(overactive bladder, treatment of; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)
- IT Coating materials  
(polymer film; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)
- IT Dissolution  
Gelation agents  
Gums and Mucilages  
Plasticizers  
(preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)
- IT Adhesins  
Agglutinins and Lectins  
Bentonite, biological studies  
Biopolymers  
Carbohydrates, biological studies  
Clays, biological studies  
Gelatins, biological studies  
Glycoproteins  
Hydrocarbon oils  
Kaolin, biological studies  
Polyesters, biological studies  
Polyoxyalkylenes, biological studies  
Smectite-group minerals  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)
- IT Drug delivery systems

(tablets, controlled-release; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Fats and Glyceridic oils, biological studies  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (vegetable, hydrogenated; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT Granulation  
 (wet; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 9003-01-4D, crosslinked  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (Carbopol; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

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 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (Crospovidone; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 9010-88-2, Eudragit NE 30D  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (Eudragit NE 50D; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 9050-36-6, Maltodextrin  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (Mor-Rex; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 7631-86-9, Silica, biological studies  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (colloidal; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 9004-34-6, Cellulose, biological studies  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (microcryst.; preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 124937-51-5, Tolterodine 124937-52-6, Tolterodine tartrate  
 RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (preparation of controlled-release polymeric mucoadhesive matrix containing tolterodine for tablets or minitablets)

IT 50-70-4, Sorbitol, biological studies 50-99-7, D-Glucose, biological studies 56-40-6, Glycine, biological studies 57-11-4, Stearic acid, biological studies 57-48-7, Fructose, biological studies 57-50-1, Sucrose, biological studies 69-65-8, D-Mannitol 77-89-4, Acetyl triethyl citrate 77-90-7, Acetyl tributyl citrate 79-41-4D, Methacrylic acid, derivs., polymers 84-66-2, Diethyl phthalate 84-74-2, Dibutyl phthalate 87-89-8, Inositol 88-99-3, Phthalic acid, biological studies 102-76-1, Triacetin 108-32-7, Propylene carbonate 109-43-3, Dibutyl sebacate 112-92-5, Stearyl alcohol 117-81-7, Diethyl phthalate 134-03-2, Sodium ascorbate 471-34-1, Calcium carbonate, biological studies 557-04-0, Magnesium stearate 557-05-1, Zinc stearate 585-86-4, Lactitol 1327-43-1, Magnesium aluminum silicate 1344-95-2, Calcium silicate 1592-23-0, Calcium stearate 4070-80-8, Sodium stearyl fumarate 7789-77-7, Dibasic calcium phosphate dihydrate 9000-01-5, Acacia gum 9000-07-1, Carrageenan 9000-11-7, Carboxymethyl cellulose 9000-30-0, Guar gum 9000-36-6, Karaya gum 9000-40-2,

Locust bean gum 9000-65-1, Tragacanth gum 9000-69-5, Pectin  
 9002-18-0, Agar 9002-88-4D, Polyethylene, alkyl ethers 9003-01-4,  
 Poly(acrylic acid) 9003-39-8, Polyvinylpyrrolidone 9004-32-4, Sodium  
 CM-cellulose 9004-38-0, CAP 9004-53-9, Dextrin 9004-57-3, Ethyl  
 cellulose 9004-61-9, Hyaluronic acid 9004-62-0, Hydroxyethyl cellulose  
 9004-64-2, Hydroxypropyl cellulose 9004-65-3, Hypromellose 9004-67-5,  
 Methyl cellulose 9005-25-8, Starch, biological studies 9005-32-7,  
 Alginic acid 9005-38-3, Sodium alginate 9005-65-6, Polysorbate 80  
 9005-82-7, Amylose 9012-76-4, Chitosan 9050-31-1, HPMCP 9063-38-1,  
 Sodium starch glycolate 10101-41-4, Calcium sulfate dihydrate  
 12705-30-5, Celutab 13463-67-7, Titanium dioxide, biological studies  
 14807-96-6, Talc, biological studies 18662-40-3, Calcium sulfate  
 monohydrate 25086-15-1, Eudragit S 100 25212-88-8, Eudragit L 100-55  
 25322-68-3, Polyethylene oxide 25496-72-4, Glyceryl monooleate  
 26009-03-0, Poly(glycolic acid) 26023-30-3, Poly[oxy(1-methyl-2-oxo-1,2-  
 ethanediyl)] 26100-51-6, Poly(DL-lactic acid) 26124-68-5,  
 Poly(glycolic acid) 26936-24-3, Eudragit FS 30D 31566-31-1, Glyceryl  
 monostearate 33434-24-1, Eudragit RS 100 36653-82-4, Cetyl alcohol  
 39301-46-7, Calcium pectinate 53237-50-6 66828-18-0, Dextrose  
 71138-97-1, HPMCAS 74811-65-7, Croscarmellose sodium 77538-19-3,  
 Glyceryl behenate 77938-63-7, Dextrose monohydrate  
 139061-06-6, Calcium lactate trihydrate 147335-38-4, Eudragit NE 40D  
 178806-61-6, Eudragit RLPO 476312-12-6, Carbopol 71G 869094-48-4,  
 Maltrons  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (preparation of controlled-release polymeric mucoadhesive matrix containing  
 tolterodine for tablets or minitablets)

L1 ANSWER 2 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:1885 CAPLUS

DOCUMENT NUMBER: 142:79974

TITLE: Soft tablet containing high molecular weight  
 cellulosics

INVENTOR(S): Wynn, David; Parikh, Nick

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 8 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2004265373	A1	20041230	US 2003-608681	20030627
CA 2472432	AA	20041227	CA 2004-2472432	20040625
EP 1498114	A1	20050119	EP 2004-253844	20040625
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR				
PRIORITY APPLN. INFO.: US 2003-607766 A 20030627				
US 2003-608681 A 20030627				

TI Soft tablet containing high molecular weight cellulosics

AB The invention relates to an immediate-release tablet capable of  
 being chewed or disintegrated in the oral cavity, which comprises an  
 active ingredient having an optional taste masking coating, and a matrix  
 comprising hydroxalkyl cellulose having a weight average mol. weight of  
 60,000-

5,000,000. The tablet has exceptionally good mouth-feel and stability. Thus, a coating solution contained cellulose acetate 43, Hypromellose phthalate 53, and Polysorbate-80 4%. Ibuprofen granules were obtained in the conventional manner and were then coated with the above taste-masking solution

ST soft tablet mol wt cellulose

IT Granulation  
 (dry granulation; soft tablet containing high mol. weight celluloses)

IT Drug delivery systems  
 (granules; soft tablet containing high mol. weight celluloses)

IT Bitterness

Coating materials

Compression

Molecular weight distribution

Viscosity  
 (soft tablet containing high mol. weight celluloses)

IT Carbohydrates, biological studies  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (soft tablet containing high mol. weight celluloses)

IT Drug delivery systems  
 (tablets, immediate release; soft tablet containing high mol. weight celluloses)

IT Drug delivery systems  
 (tablets; soft tablet containing high mol. weight celluloses)

IT 9004-34-6, Cellulose, biological studies  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (microcryst.; soft tablet containing high mol. weight celluloses)

IT 50-70-4, Sorbitol, biological studies 50-78-2, Acetylsalicylic acid  
 58-73-1, Diphenhydramine 69-65-8, Mannitol 87-99-0, Xylitol 90-82-4,  
 Pseudoephedrine 103-90-2, Acetaminophen 125-71-3, Dextromethorphan  
 132-22-9, Chlorpheniramine 303-53-7, Cyclobenzaprine 5104-49-4,  
 Flurbiprofen 9004-34-6D, Cellulose, ethers 9004-35-7 9004-62-0,  
 Hydroxyethyl cellulose 9004-64-2, Hydroxypropyl cellulose 9004-65-3,  
 Hydroxypropyl methyl cellulose 9032-42-2, Hydroxyethyl methyl cellulose  
 9050-31-1, Hypromellose phthalate 14838-15-4, Phenylpropanolamine  
 15307-86-5, Diclofenac 15687-27-1, Ibuprofen 22071-15-4, Ketoprofen  
 22204-53-1, Naproxen 37353-59-6, Hydroxymethyl cellulose 50679-08-8,  
 Terfenadine 68844-77-9, Astemizole 71125-38-7, Meloxicam 77938-63-7,  
 Dextrose monohydrate 79794-75-5, Loratadine  
 83799-24-0, Fexofenadine 83881-51-0, Cetirizine 162011-90-7, Rofecoxib  
 169590-42-5, Celecoxib  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (soft tablet containing high mol. weight celluloses)

L1 ANSWER 3 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 2005:993 CAPLUS  
 DOCUMENT NUMBER: 142:79963  
 TITLE: Soft tablets containing high molecular weight celluloses  
 INVENTOR(S): Wynn, David; Parikh, Nick  
 PATENT ASSIGNEE(S): USA  
 SOURCE: U.S. Pat. Appl. Publ., 9 pp.  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

## PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2004265372	A1	20041230	US 2003-607766	20030627
CA 2472432	AA	20041227	CA 2004-2472432	20040625
EP 1491184	A1	20041229	EP 2004-253843	20040625
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR				
PRIORITY APPLN. INFO.:			US 2003-607766	A 20030627
			US 2003-608681	A 20030627

- TI Soft tablets containing high molecular weight celluloses  
 AB An immediate release tablet capable of being chewed or subjected to disintegration in the oral cavity, comprises an active ingredient having an optional taste-masking coating, and a matrix comprising hydroxyalkyl cellulose having a weight average mol. weight of 60,000-5,000,000. The tablet has exceptionally good mouth-feel and stability. A coating solution was prepared by dispersing cellulose acetate 43, Hypromellose phthalate 53, and Polysorbate-80 4% in a solvent consisting of 90% acetone and 10% water under ambient conditions, so that the finished solution contained 10% of the coating materials. Ibuprofen granules prepared in the conventional way were then coated with the above taste-masking solution. High weight average mol. weight hydroxyalkyl cellulose-containing tablets had significantly less of a grittiness feel in the mouth in comparison to those tablets lacking the high weight average mol. weight hydroxyalkyl cellulose.  
 ST soft tablet mol wt cellulose  
 IT Granulation  
     (dry granulation; soft tablets containing high mol. weight celluloses)  
 IT Bitterness  
 Coating materials  
 Compression  
 Dissolution  
 Molecular weight distribution  
 Solubilizers  
     (soft tablets containing high mol. weight celluloses)  
 IT Carbohydrates, biological studies  
 Polymers, biological studies  
 Polyoxyalkylenes, biological studies  
 Shellac  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
     (soft tablets containing high mol. weight celluloses)  
 IT Drug delivery systems  
     (tablets, enteric-coated; soft tablets containing high mol. weight celluloses)  
 IT Drug delivery systems  
     (tablets, immediate release; soft tablets containing high mol. weight celluloses)  
 IT 50-70-4, Sorbitol, biological studies 50-78-2, Acetylsalicylic acid, 58-73-1, Diphenhydramine 69-65-8, Mannitol 79-41-4D, Methacrylic acid, esters, polymers 87-99-0, Xylitol 90-82-4, Pseudoephedrine 103-90-2, Acetaminophen 125-71-3, Dextromethorphan 132-22-9, Chlorpheniramine 303-53-7, Cyclobenzaprine 5104-49-4, Flurbiprofen 9002-89-5, Poly(vinyl alcohol) 9003-39-8, Polyvinylpyrrolidone 9004-32-4

9004-34-6D, Cellulose, ethers 9004-35-7 9004-36-8, Cellulose acetate butyrate 9004-38-0, Cellulose acetate phthalate 9004-57-3, Ethyl cellulose 9004-62-0, Hydroxyethyl cellulose 9004-64-2, Hydroxypropyl cellulose 9004-65-3, Hydroxypropyl methyl cellulose 9004-67-5, Methyl cellulose 9010-88-2, Ethyl acrylate-methyl methacrylate copolymer 9012-09-3, Cellulose triacetate 9032-42-2, Hydroxyethyl methyl cellulose 9050-31-1, Hydroxypropyl methyl cellulose phthalate 14838-15-4, Phenylpropanolamine 15307-86-5, Diclofenac 15687-27-1, Ibuprofen 22071-15-4, Ketoprofen 22204-53-1, Naproxen 25322-68-3, Polyethylene glycol 37353-59-6, Hydroxymethyl cellulose 50679-08-8, Terfenadine 53237-50-6, Polyvinyl acetate phthalate 68844-77-9, Astemizole 70535-77-2, Hydroxypropyl methyl cellulose acetate succinate 71125-38-7, Meloxicam 77938-63-7, Dextrose monohydrate 79794-75-5, Loratadine 83799-24-0, Fexofenadine 83881-51-0, Cetirizine 162011-90-7, Rofecoxib 169590-42-5, Celecoxib  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(soft tablets containing high mol. weight celluloses)

L1 ANSWER 4 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:368929 CAPLUS  
DOCUMENT NUMBER: 140:363062  
TITLE: Pharmaceutical compositions of ganciclovir  
INVENTOR(S): Mathur, Rajeev Shankar; Kumar, Pananchukunnath Manoj; Roy, Sunilendu Bhushan; Malik, Rajiv  
PATENT ASSIGNEE(S): Ranbaxy Laboratories Limited, India  
SOURCE: PCT Int. Appl., 21 pp.  
CODEN: PIXXD2  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004037263	A1	20040506	WO 2003-IB4664	20031022
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
AU 2003274410	A1	20040513	AU 2003-274410	20031022
EP 1556050	A1	20050727	EP 2003-758391	20031022
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK				
US 2006189565	A1	20060824	US 2006-532024	20060407
PRIORITY APPLN. INFO.:			IN 2002-DE1058	A 20021022
			WO 2003-IB4664	W 20031022
REFERENCE COUNT:	4	THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT		
IT Drug delivery systems (tablets; stable pharmaceutical compns. of ganciclovir)				
IT 50-70-4, Sorbitol, biological studies	50-99-7, Glucose, biological			

studies 57-50-1, Sucrose, biological studies 63-42-3, Lactose 69-65-8, D-Mannitol 7789-77-7, Dibasic calcium phosphate dihydrate 9000-01-5, Acacia gum 9000-30-0, Guar gum 9000-65-1, Tragant gum 9003-39-8, PVP 9004-32-4, Sodium CMC 9004-64-2, Hydroxypropyl cellulose 9004-65-3, Hydroxypropyl methyl cellulose 9004-67-5, Methyl cellulose 9005-25-8, Starch, biological studies 9005-32-7, Alginic acid 9063-38-1, Sodium starch glycolate 10031-30-8 10101-41-4, Calcium sulfate dihydrate 25322-68-3, Polyethylene glycol 74811-65-7, Croscarmellose sodium 77938-63-7, Dextrose monohydrate 82410-32-0, Ganciclovir 139061-06-6  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (stable pharmaceutical compns. of ganciclovir)

L1 ANSWER 5 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 2004:145843 CAPLUS  
 DOCUMENT NUMBER: 141:355096  
 TITLE: Powdered and granular materials used in the fabrication of compressed tablets  
 AUTHOR(S): Delattre, Luc  
 CORPORATE SOURCE: Laboratoire de Technologie Pharmaceutique, Departement de Pharmacie, Faculte de Medecine, Universite de Liege, Liege, Belg.  
 SOURCE: Bulletin de la Societe Royale des Sciences de Liege (2003), 72(5), 317-339  
 CODEN: BSRSA6; ISSN: 0037-9565  
 PUBLISHER: Societe Royale des Sciences de Liege  
 DOCUMENT TYPE: Journal  
 LANGUAGE: French  
 REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

TI Powdered and granular materials used in the fabrication of compressed tablets  
 AB The effect of Mg stearate mixing time on the crushing strength of tablets was determined  
 ST compressed tablet property  
 IT Drug delivery systems  
     (granules; powdered and granular materials in fabrication of compressed tablets)  
 IT Compaction  
 Compression  
 Crushing strength  
 Shear  
     (powdered and granular materials in fabrication of compressed tablets)  
 IT Drug delivery systems  
     (tablets; powdered and granular materials in fabrication of compressed tablets)  
 IT Granulation  
     (wet; powdered and granular materials in fabrication of compressed tablets)  
 IT 9004-34-6, Avicel PH102, biological studies  
 RL: PEP (Physical, engineering or chemical process); PYP (Physical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)  
     (microcryst.; powdered and granular materials in fabrication of compressed tablets)  
 IT 63-42-3, Tablettose 557-04-0 5965-66-2, Pharmatose DCL 21 7789-77-7,

Dibasic calcium phosphate dihydrate 12705-30-5, Celutab 64044-51-5,  
 Lactose monohydrate 77938-63-7, Dextrose monohydrate  
 RL: PEP (Physical, engineering or chemical process); PYP (Physical  
 process); THU (Therapeutic use); BIOL (Biological study); PROC (Process);  
 USES (Uses)  
 (powdered and granular materials in fabrication of compressed  
 tablets)

L1 ANSWER 6 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:737151 CAPLUS  
 DOCUMENT NUMBER: 139:250306  
 TITLE: Soft tablets containing high molecular  
 weight polyethylene oxide  
 INVENTOR(S): Luber, Joseph; Bunick, Frank J.  
 PATENT ASSIGNEE(S): McNeil-PPC, Inc., USA  
 SOURCE: U.S. Pat. Appl. Publ., 7 pp.  
 CODEN: USXXCO  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2003175336	A1	20030918	US 2002-97000	20020313
US 6753009	B2	20040622		
CA 2421685	AA	20030913	CA 2003-2421685 US 2002-97000	20030312 A 20020313

PRIORITY APPLN. INFO.:  
 REFERENCE COUNT: 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS  
 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

TI Soft tablets containing high molecular weight polyethylene oxide  
 AB The invention relates to an immediate release tablet capable of  
 being chewed or disintegrated in the oral cavity, which comprises a  
 pharmaceutically active ingredient, and a matrix comprising polyethylene  
 oxide having a weight average mol. weight of from about 500,000 to about  
 10,000,000.

The tablet possesses exceptionally good mouthfeel and stability.  
 For example, tablets were formulated containing polyethylene oxide  
 (average mol. weight 5,000,000), vitamin E granules 13.3, erythritol 100,  
 crospovidone 25, colorant 2.5, coated ibuprofen 282.1, flavors 15,  
 sucralose 10, dextrose monohydrate 658, and lubricants  
 7.5 parts.

ST immediate release soft tablet matrix PEG

IT Antacids

Antioxidants

(immediate-release matrixes containing high mol. weight PEG and antioxidants  
 for soft tablets)

IT Polyoxyalkylenes, biological studies

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(immediate-release matrixes containing high mol. weight PEG and antioxidants  
 for soft tablets)

IT Drug delivery systems

(tablets, buccal; immediate-release matrixes containing high mol.  
 weight PEG and antioxidants for soft tablets)

IT Drug delivery systems

(tablets, chewable; immediate-release matrixes containing high  
 mol. weight PEG and antioxidants for soft tablets)

IT Drug delivery systems  
     (tablets, controlled-release; immediate-release matrixes  
     containing high mol. weight PEG and antioxidants for soft tablets)

IT 50-78-2, Acetylsalicylic acid 58-73-1, Diphenhydramine 59-02-9,  
   α-Tocopherol 90-82-4, Pseudoephedrine 103-90-2, Acetaminophen  
   113-92-8 121-79-9, Propyl gallate 125-71-3, Dextromethorphan  
   128-37-0, biological studies 303-53-7, Cyclobenzaprine 319-89-1,  
   Tetrahydroxyquinone 603-50-9, Bisacodyl 915-30-0, Diphenoxylate  
   1406-18-4, Vitamin E 5104-49-4, Flurbiprofen 7397-62-8, Butyl  
   hydroxyacetate 7440-69-9, Bismuth, biological studies 9031-11-2,  
   Lactase 14838-15-4, Phenylpropanolamine 15307-86-5, Diclofenac  
   15687-27-1, Ibuprofen 22071-15-4, Ketoprofen 22204-53-1, Naproxen  
   25322-68-3, Polyethylene oxide 50679-08-8, Terfenadine 51481-61-9,  
   Cimetidine 53179-11-6, Loperamide 66357-35-5, Ranitidine 68844-77-9,  
   Astemizole 71125-38-7, Meloxicam 76824-35-6, Famotidine 79794-75-5,  
   Loratadine 83799-24-0, Fexofenadine 83881-51-0, Cetirizine  
   162011-90-7, Rofecoxib 169590-42-5, Celecoxib 179474-81-8,  
   Prucalopride  
   RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
     (immediate-release matrixes containing high mol. weight PEG and antioxidants  
     for soft tablets)

L1 ANSWER 7 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:8105 CAPLUS  
 DOCUMENT NUMBER: 138:61356  
 TITLE: Method to aid smoking cessation using dextrose and/or levulose  
 INVENTOR(S): West, Robert; Hajek, Peter  
 PATENT ASSIGNEE(S): UK  
 SOURCE: Brit. UK Pat. Appl., 7 pp.  
 CODEN: BAXXDU  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
GB 2376885	A1	20021231	GB 2001-15568	20010626
PRIORITY APPLN. INFO.:			GB 2001-15568	20010626
AB	The present invention concerns methods of treating patients for nicotine and tobacco addiction, for alleviating nicotine withdrawal, for improving the effects of other smoking cessation therapies and as longterm smoking cessation maintenance therapy. The invention comprises pharmaceutical compns. comprising dextrose monohydrate and/or levulose in combination with amfebutamone or any other non-nicotine smoking cessation method whose efficacy can be enhanced by addition of dextrose or levulose. Specific combinations of drugs (dextrose and/or levulose combined with amfebutamone) as well as dextrose and/or levulose in combination with certain drug classes (e.g., stimulant drugs, antidepressants, and drugs used in treatment of psychoactive substance use disorders) are described. These compns. are also contemplated for use in the treatment of alcoholism, cocaine dependence and other drug dependencies.			
IT	Drug delivery systems (tablets, chewable; compns. containing dextrose and/or levulose in combination with amfebutamone for smoking cessation and treatment of			

alcoholism and drug dependence)

L1 ANSWER 8 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 2002:674574 CAPLUS  
 DOCUMENT NUMBER: 137:206555  
 TITLE: Soft tablet containing dextrose monohydrate  
 INVENTOR(S): Bunick, Frank J.; Luber, Joseph  
 PATENT ASSIGNEE(S): USA  
 SOURCE: U.S. Pat. Appl. Publ., 5 pp.  
 CODEN: USXXCO  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2002122823	A1	20020905	US 2000-752899 US 2000-752899	20001229 20001229
PRIORITY APPLN. INFO.:				
TI Soft tablet containing dextrose monohydrate				
AB A tablet capable of being chewed or disintegrated in the oral cavity, comprises an active ingredient, and a matrix containing directly compressible dextrose monohydrate and sucralose, the tablet being substantially fat free and the matrix being substantially free of non-saccharide water-soluble polymeric binders. Thus, tablets contained sucralose 8.0 FD&C Yellow #6 Al Lake 3.0, orange flavor 10.0 Crospovidone 15.0, coated ibuprofen 140.6, dextrose monohydrate 850.0, and Mg stearate 7.5 mg/tablet.				
ST soft tablet dextrose monohydrate				
IT Antioxidants				
Compression				
Dyes				
Flavoring materials				
Granulation				
Human				
Lubricants				
Particle size distribution				
Preservatives				
Surfactants				
Sweetening agents (soft tablets containing dextrose monohydrate )				
IT Drug delivery systems (tablets; soft tablets containing dextrose monohydrate)				
IT 9003-39-8, Polyvinylpyrrolidone 9004-32-4, Carboxymethyl cellulose RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (crosslinked; soft tablets containing dextrose monohydrate)				
IT 9004-34-6, Cellulose, biological studies RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (microcryst.; soft tablets containing dextrose monohydrate)				
IT 57-11-4, Stearic acid, biological studies 58-73-1, Diphenhydramine 90-82-4, Pseudoephedrine 103-90-2, Acetaminophen 113-92-8, Chlorpheniramine 125-71-3, Dextromethorphan 471-34-1, Calcium				

carbonate, biological studies 546-93-0, Magnesium carbonate 557-04-0, Magnesium stearate 1309-42-8, Magnesium hydroxide 1309-48-4, Magnesium oxide, biological studies 2783-94-0, FD&C Yellow #6 9005-25-8, Starch, biological studies 9063-38-1, Sodium starch glycolate 14431-43-7, Dextrose monohydrate 15687-27-1, Ibuprofen 21645-51-2, Aluminum hydroxide, biological studies 56038-13-2, Sucralose RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (soft tablets containing dextrose monohydrate )

L1 ANSWER 9 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2002:674573 CAPLUS

DOCUMENT NUMBER: 137:206554

TITLE: Chewable tablets containing hydrate excipients.

INVENTOR(S): Bunick, Frank J.; Luber, Joseph

PATENT ASSIGNEE(S): McNeil-PPC, Inc., USA

SOURCE: U.S. Pat. Appl. Publ., 5 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2002122822	A1	20020905	US 2000-752601	20001229
US 6814978	B2	20041109		
US 2003175339	A1	20030918	US 2003-413804	20030415
PRIORITY APPLN. INFO.:			US 2000-752601	A1 20001229
REFERENCE COUNT:	18	THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT		

TI Chewable tablets containing hydrate excipients.

AB The invention relates to a process for preparing a soft tablet capable of being chewed or disintegrated in the oral cavity. The tablet is prepared by forming a tablet having a friability of less than about 2% from a mixture comprising a pharmaceutically active ingredient, an excipient in the form of a hydrate, and a water-swellable excipient, and then applying sufficient energy, preferably in the form of heat, to the tablet for a sufficient time to decrease the hardness of the tablet by at least about 20%. A composition contained sucralose 8.0, coated ibuprofen (69.0%) 140.6, flavor 10.0, dextrose monohydrate 850.0, Crospovidone 15.0, and Mg stearate 7.5.

ST tablet chewable hydrate excipient

IT Compression

Härdföring (mechanical)

Particle size

(chewable tablets containing hydrate excipients)

IT Drug delivery systems

(tablets, chewable; chewable tablets containing hydrate excipients)

IT 9003-39-8D, crosslinked

RL: MOA (Modifier or additive use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(Crospovidone; chewable tablets containing hydrate excipients)

IT 5949-29-1, Citric acid monohydrate 7782-85-6, Phosphoric acid, disodium

salt, heptahydrate 7789-77-7, Dibasic calcium phosphate dihydrate 9004-34-6, Cellulose, biological studies 9004-53-9, Dextrin 9005-25-8, Starch, biological studies 9005-32-7, Alginic acid 9050-36-6, Maltodextrin 9063-38-1, Sodium starch glycolate 10028-24-7, Phosphoric acid, disodium salt, dihydrate 10039-32-4, Phosphoric acid, disodium salt, dodecahydrate 10049-21-5, Monosodium phosphate monohydrate 13472-35-0, Monosodium phosphate dihydrate 14431-43-7, Dextrose monohydrate 64044-51-5, Lactose monohydrate 74811-65-7, Croscarmellose sodium

RL: MOA (Modifier or additive use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(chewable tablets containing hydrate excipients)

IT 58-73-1, Diphenhydramine 90-82-4, Pseudoephedrine 103-90-2, Acetaminophen 113-92-8, Chlorpheniramine 125-71-3, Dextromethorphan 471-34-1, Calcium carbonate, biological studies 546-93-0, Magnesium carbonate 1309-42-8, Magnesium hydroxide 1309-48-4, Magnesium oxide, biological studies 15687-27-1, Ibuprofen 21645-51-2, Aluminum hydroxide, biological studies

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(chewable tablets containing hydrate excipients)

L1 ANSWER 10 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:434870 CAPLUS

DOCUMENT NUMBER: 135:51047

TITLE: Nanoparticulate eplerenone compositions

INVENTOR(S): Thosar, Shilpa S.; Gokhale, Rajeev D.; Tolbert, Dwain S.; Desai, Subhash

PATENT ASSIGNEE(S): Pharmacia Corporation, USA

SOURCE: PCT Int. Appl., 64 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001041770	A2	20010614	WO 2000-US30179	20001204
WO 2001041770	A3	20011122		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
AU 2001017562	A5	20010618	AU 2001-17562	20001204
EP 1175220	A2	20020130	EP 2000-980277	20001204
EP 1175220	B1	20050427		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO				
EP 1527782	A1	20050504	EP 2004-30120	20001204
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI, CY, TR				
AT 293977	E	20050515	AT 2000-980277	20001204

PT 1175220	T	20050729	PT 2000-980277	20001204
ES 2240209	T3	20051016	ES 2000-980277	20001204
US 2002006919	A1	20020117	US 2000-732246	20001207
US 2003212053	A1	20031113	US 2003-417602	20030416
PRIORITY APPLN. INFO.:				
			US 1999-169658P	P 19991208
			US 2000-208981P	P 20000602
			EP 2000-980277	A3 20001204
			WO 2000-US30179	W 20001204
			US 2000-732246	A3 20001207

AB There is provided a pharmaceutical composition comprising eplerenone in solid particulate form, wherein at least 90 of the eplerenone particles are smaller than about 15  $\mu\text{m}$ , for example about 0.01 to about 1  $\mu\text{m}$ , in diameter. The composition can be adapted for oral administration, for example

as a

tablet or capsule comprising eplerenone in a unit dosage amount of about 10 to about 1000 mg and one or more excipients. An immediate release tablet was prepared containing nanoparticulate eplerenone 25.00, lactose monohydrate 35.70, microcryst. cellulose 15.38, croscarmellose sodium 4.25, HPMC 2.55, Na lauryl sulfate 0.85, Mg stearate 0.42, and Opadry White YS-1-18027A 2.55 mg/tablet.

ST eplerenone nanoparticle tablet capsule

IT Drug delivery systems

(tablets; nanoparticulate eplerenone compns.)

IT 50-70-4, Sorbitol, biological studies 50-99-7, Dextrose, biological studies 56-40-6, Glycine, biological studies 57-11-4, Stearic acid, biological studies 57-50-1, Sucrose, biological studies 63-42-3, Lactose 69-65-8, D-Mannitol 87-89-8, Inositol 87-99-0, Xylitol 112-80-1, Oleic acid, biological studies 121-54-0, Benzethonium chloride 123-03-5, Cetylpyridinium chloride 127-09-3, Sodium acetate 143-19-1, Sodium oleate 151-21-3, Sodium lauryl sulfate, biological studies 328-39-2, Leucine 471-34-1, Calcium carbonate, biological studies 532-32-1, Sodium benzoate 557-04-0, Magnesium stearate 577-11-7, Dioctyl sodium sulfosuccinate 822-16-2, Sodium stearate 1327-43-1, Magnesium aluminum silicate 1338-39-2, Sorbitan monolaurate 1338-41-6, Sorbitan monostearate 1338-43-8, Sorbitan monooleate 1592-23-0, Calcium stearate 2717-15-9, Triethanolamine oleate 3097-08-3, Magnesium lauryl sulfate 7631-86-9, Silica, biological studies 7647-14-5, Sodium chloride, biological studies 7704-73-6, Sodium fumarate 7789-77-7, Dicalcium phosphate dihydrate 9000-30-0, Guar gum 9000-36-6, Karaya gum 9000-40-2, Locust bean gum 9000-65-1, Gum tragacanth 9000-69-5, Pectin 9002-18-0, Agar 9003-39-8, Pvp 9004-32-4 9004-32-4, CM-cellulose 9004-34-6, Cellulose, biological studies 9004-57-3, Ethyl cellulose 9004-64-2, Hydroxypropyl cellulose 9004-65-3, HPMC 9004-67-5, Methyl cellulose 9004-99-3, Polyoxyethylene stearate 9005-25-8, Starch, biological studies 9005-32-7, Alginic acid 9005-64-5, Polysorbate 20 9005-65-6, Polysorbate 80 9005-82-7, Amylose 9036-19-5, Octoxynol 9 9063-38-1, Sodium starch glycolate 10043-35-3, Boric acid, biological studies 10101-41-4, Calcium sulfate dihydrate 14431-43-7, Dextrose monohydrate 14807-96-6, Talc, biological studies 18641-57-1, Glyceryl behenate 18662-40-3, Sulfuric acid, calcium salt (1:1), monohydrate 25301-02-4, Tyloxapol 25322-68-3, Peg 26027-38-3, Nonoxynol 9 26266-57-9, Sorbitan monopalmitate 27306-76-9, Polyoxyethylene cetylstearyl ether 31566-31-1, Glyceryl monostearate 37321-62-3, Propylene glycol laurate 64044-51-5, Lactose monohydrate 66828-18-0, Dextrose 74811-65-7, Croscarmellose sodium 106392-12-5, Poloxamer 139061-06-6, Propanoic acid, 2-hydroxy-, calcium salt (2:1), trihydrate

RL: MOA (Modifier or additive use); THU (Therapeutic use); BIOL  
 (Biological study); USES (Uses)  
 (nanoparticulate eplerenone compns.)

L1 ANSWER 11 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 2001:136959 CAPLUS  
 DOCUMENT NUMBER: 134:183494  
 TITLE: Orally dissolvable prenatal multi-vitamin  
 INVENTOR(S): Devries, Tina; Valentine, William; Valentine, William K.  
 PATENT ASSIGNEE(S): Warner Chilcott Laboratories Ireland Limited, USA  
 SOURCE: PCT Int. Appl., 38 pp.  
 CODEN: PIXXD2  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 2  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001011991	A1	20010222	WO 2000-US40557	20000803
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
US 6495177	B1	20021217	US 2000-539850	20000331
PRIORITY APPLN. INFO.:			US 1999-148803P	P 19990813
			US 1999-148806P	P 19990813
			US 2000-539850	A 20000331

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

AB The present invention provides an orally administrable nutritional supplement which is highly palatable, such as a chewable prenatal vitamin/mineral supplement. The supplement is preferably made in the form of a tablet that, upon chewing, dissolves rapidly in the mouth. The tablet is particularly suitable for administration of vitamins and minerals to women during pregnancy. The invention also includes methods of making and using such supplements.

ST vitamin mineral supplement tablet

IT Drug delivery systems

(tablets, chewable; orally dissolvable prenatal multi-vitamin)

IT 50-70-4, Sorbitol, biological studies 50-81-7, Vitamin C, biological studies 50-99-7, Dextrose, biological studies 57-48-7, D-Fructose, biological studies 57-50-1, Sucrose, biological studies 58-86-6, D-Xylose, biological studies 58-95-7, Vitamin E acetate 59-30-3, Folic acid, biological studies 59-30-3D, Folic acid, salts 59-43-8, Vitamin B1, biological studies 59-67-6, Niacin, biological studies 63-42-3, Lactose 67-97-0, Vitamin D3 68-19-9, Vitamin B12 69-65-8, Mannitol 69-79-4, Maltose 83-88-5, Vitamin B2, biological studies 98-92-0, Niacinamide 134-03-2, Sodium ascorbate 141-01-5, Ferrous fumarate 557-04-0, Magnesium stearate 1406-18-4, Vitamin E 7235-40-7,

$\beta$ -Carotene 7439-89-6, Iron, biological studies 7439-89-6D, Iron, compds., biological studies 7440-70-2, Calcium, biological studies 7758-87-4, Tricalcium phosphate 8059-24-3, Vitamin B6 9003-39-8, Polyvinyl pyrrolidone 9004-34-6, Cellulose, biological studies 9016-00-6, Dimethyl polysiloxane 9050-36-6, Maltodextrin 11103-57-4, Provitamin A 14431-43-7, Dextrose monohydrate  
 RL: FFD (Food or feed use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (orally dissolvable prenatal multi-vitamin)

L1 ANSWER 12 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1997:471006 CAPLUS

DOCUMENT NUMBER: 127:152888

TITLE: Potassium carbonate as a desiccant in effervescent tablets

AUTHOR(S): Wells, Mickey L.; Wood, Daniel L.; Sanftleben, Ronald; Shaw, Kelley; Hottovy, Jeff; Weber, Thomas; Geoffroy, Jean-Marie; Alkire, Todd G.; Emptage, Michael R.; Sarabia, Rafael

CORPORATE SOURCE: Glaxo Wellcome Inc., Research Triangle Park, NC, 27709, USA

SOURCE: International Journal of Pharmaceutics (1997), 152(2), 227-235

CODEN: IJPHDE; ISSN: 0378-5173

PUBLISHER: Elsevier

DOCUMENT TYPE: Journal

LANGUAGE: English

TI Potassium carbonate as a desiccant in effervescent tablets

AB A central composite study design was used to determine the moisture scavenging effect of 0-2% weight/weight potassium carbonate in an effervescent dosage form containing 0.2-1.3% weight/weight total moisture. Total moisture content of the

tablets was calculated by adding the water contribution of each ingredient based on loss on drying or Karl Fischer data. Tablets were directly compressed on a rotary tablet press, packaged in cold form foil/foil blisters, and then thermally stressed by exposure to 75°C for 3 h. In this study, exposure of effervescence in such a manner has been shown to release water of hydration from dextrose monohydrate, thus giving a convenient means of adding water and then 'activating' it to perform rapid moisture stability studies. After thermal stressing, tablets were given a rating from 0-7 (least to most) as to the degree of tablet mottling due to effervescent base degradation. Response surface regression of the data resulted in a quadratic equation with an adjusted R<sup>2</sup> of 0.92 and no evidence of lack of fit ( $P = 0.85$ ). Anal. of the data showed the optimal level of potassium carbonate to be 1.3% weight/weight for the formulations tested. This level of potassium carbonate will accommodate total moisture levels up to 0.4% weight/weight and still prevent effervescent base degradation.

ST potassium carbonate desiccant effervescent pharmaceutical tablet

IT Particle size

(potassium carbonate as desiccant in effervescent tablets)

IT Drying agents

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(potassium carbonate as desiccant in effervescent tablets)

IT Drug delivery systems

Drug delivery systems

(tablets, effervescent; potassium carbonate as desiccant in

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effervescent tablets)  
IT 584-08-7, Potassium carbonate  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(potassium carbonate as desiccant in effervescent tablets)

L1 ANSWER 13 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1997:204347 CAPLUS  
DOCUMENT NUMBER: 126:255506  
TITLE: Compressed tablet transitory lubricant system  
INVENTOR(S): Valentine, William; Valentine, William K.  
PATENT ASSIGNEE(S): Advanced Technology Pharmaceuticals Corporation, USA  
SOURCE: U.S., 6 pp.  
CODEN: USXXAM  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5609883	A	19970311	US 1994-307922	19940916
PRIORITY APPLN. INFO.:			US 1994-307922	19940916

TI Compressed tablet transitory lubricant system  
AB A method is provided for making fast dissolving storage stable tablets by compression on standard high speed tablet production machinery wherein the formulation contains a carbohydrate having a special particle size and/or structure, in combination with controlled amts. of a transitory liquid as a lubricant, which liquid is removed following compression. Dextrose monohydrate/maltodextrin coagglomerate 845.6, 33.3% coated chlorpheniramine maleate 3.4, 33.3% coated pseudoephedrine.HCl 50.0, 10% dextromethorphan.HBr magnesium trisilicate 56.0, spray dried lemon flavor 45.0 g, and Et alc. 44 mL were blended then ethanol was added and mixed until a uniformly damp granulation was formed. The damp granulation was pressed on a tablet press and dried at 37° for 30 min. The finished tablets increased in hardness to 5-6 Kp and demonstrated enhanced liquefactive characteristics.  
ST compressed pharmaceutical tablet carbohydrate ethanol  
IT Lubricants  
Particle size  
(fast dissolving compressed tablet with enhanced liquefactive character)  
IT Alcohols, uses  
RL: NUU (Other use, unclassified); USES (Uses)  
(fast dissolving compressed tablet with enhanced liquefactive character)  
IT Carbohydrates, biological studies  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(fast dissolving compressed tablet with enhanced liquefactive character)  
IT Drug delivery systems  
(tablets, compressed; fast dissolving compressed tablet with enhanced liquefactive character)  
IT 64-17-5, Ethanol, uses  
RL: NUU (Other use, unclassified); USES (Uses)  
(fast dissolving compressed tablet with enhanced liquefactive

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character)  
IT 50-99-7, Dextrose, biological studies 113-92-8, Chlorpheniramine maleate  
125-69-9, Dextromethorphan hydrobromide 345-78-8, Pseudoephedrine  
hydrochloride 9050-36-6, Maltodextrin 14431-43-7, Dextrose  
monohydrate  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(fast dissolving compressed tablet with enhanced liquefactive  
character)

L1 ANSWER 14 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1996:155953 CAPLUS  
DOCUMENT NUMBER: 124:270037  
TITLE: Using starch in tabletting  
AUTHOR(S): Vanhemelrijk, J.; Heume, M.  
CORPORATE SOURCE: Cerestar Euro Centre Food, Vilvoorde, 1800, Belg.  
SOURCE: Agro-Food-Industry Hi-Tech (1995), 6(5), 9-10  
CODEN: AIHTEI; ISSN: 1120-6012  
PUBLISHER: TeknoScienze srl  
DOCUMENT TYPE: Journal; General Review  
LANGUAGE: English

AB A review with no refs. Starch, in its many basic and chemical or phys. modified forms has been used for many years in tablet production. Its hydrolysis products, maltodextrin, glucose syrup solids and dextrose monohydrate all find specialist performance niches. In addition, dextrose when fully hydrogenated to sorbitol offers a newer tabletting agent with specialist potential. Work being carried out on tabletting with starch and derivs. is described.

ST review starch tablet  
IT Pharmaceutical dosage forms  
(tablets, starch in tabletting)

L1 ANSWER 15 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1995:694438 CAPLUS  
DOCUMENT NUMBER: 123:93108  
TITLE: Effect of different excipients on release characteristics of acetylsalicylic acid from compressed pellets  
AUTHOR(S): Torrado-Santiago; Torrado, J. J.; Cadorniga, R.  
CORPORATE SOURCE: Fac. Pharm., Complutense Univ., Madrid, Spain  
SOURCE: Pharmazie (1995), 50(7), 476-8  
CODEN: PHARAT; ISSN: 0031-7144  
PUBLISHER: Gova-Verlag Pharmazeutischer Verlag  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB The release of acetylsalicylic acid matrix tablets prepared from pellets was studied with different hydrophilic excipients [microcryst. cellulose (Avicel PH 101), wheat starch and dextrose monohydrate] in different proportions. The release process was zero-order or first-order. The dissoln. efficiency varied between 23 and 75% in 8 h. MCC is the excipient with a higher compression protecting effect on the pellets during tablet compaction. In vitro drug release depends on the MCC content of the tablets.

L1 ANSWER 16 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1993:678520 CAPLUS  
DOCUMENT NUMBER: 119:278520  
TITLE: "In vitro" drug release of AAS matrix tablets

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AUTHOR(S): Torrado, S.; Torrado, Susana; Torrado V., J.; Cadorniga, R.  
CORPORATE SOURCE: Univ. Complutense, Madrid, 28040, Spain  
SOURCE: Proc. Int. Symp. Controlled Release Bioact. Mater., 20th (1993), 370-1. Editor(s): Roseman, Theodore J.; Peppas, Nicholas A.; Gabelnick, Henry L. Controlled Release Soc.: Deerfield, Ill.  
CODEN: 59LOAL  
DOCUMENT TYPE: Conference  
LANGUAGE: English  
TI "In vitro" drug release of AAS matrix tablets  
AB Matrix tablets of acetylsalicylic acid (AAS) were produced by compression of AAS coated pellet with acrylic resins (Eudragit RS). The drug release profile of the AAS pellets after compression with different excipients (microcryst. cellulose, starch and dextrose monohydrate) was studied.  
ST acetylsalicylate release matrix tablet  
IT Solution rate  
    (of acetylsalicylic acid, from matrix tablets)  
IT Pharmaceutical dosage forms  
    (tablets, matrix, acetylsalicylic acid release from)  
IT 50-78-2, Acetylsalicylic acid  
RL: PROC (Process)  
    (release of, from matrix tablets)  
  
L1 ANSWER 17 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1988:479708 CAPLUS  
DOCUMENT NUMBER: 109:79708  
TITLE: Sustained-release pharmaceutical containing fatty acid sugar esters as excipients  
INVENTOR(S): Jansen, Frans Herwigjan; Hendrickx, Jean  
PATENT ASSIGNEE(S): Sanico, N. V., Belg.; N. V. Gantax S. A.  
SOURCE: Eur. Pat. Appl., 13 pp.  
CODEN: EPXXDW  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 230332 R: AT, BE, CH, NL 8600050 FI 8700092 NO 8700104 DK 8700161 ZA 8700216 JP 62209025	A1 DE, ES, FR, GB, GR, IT, LI, LU, NL, SE A A A A A A2	19870729 19870803 19870714 19870714 19870714 19870826 19870914	EP 1987-200031 NL 1986-50 FI 1987-92 NO 1987-104 DK 1987-161 ZA 1987-216 JP 1987-7403 NL 1986-50	19870112 19860113 19870112 19870112 19870113 19870113 19870113 A 19860113

PRIORITY APPLN. INFO.: NL 1986-50 A 19860113  
AB A sustained-release pharmaceutical composition, especially in tablet form, comprises an active component, a C10-15 fatty acid sugar ester, and other appropriate substances. A tablet composition containing ibuprofen (I) 400, dextrose monohydrate 60, polyvidone 18, sucrose monopalmitate 100, stearic acid 1, talc 16, and Mg stearate 5 kg was prepared and pressed into 500,000 tablets giving 24-h release of I. Serum release of an 800 mg dose of I was 2 µg/mL initially and 18

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ST  $\mu\text{g/mL}$  after 10 h (peak), and 2  $\mu\text{g/mL}$  after 24 h.  
ibuprofen controlled release tablet sucrose ester

IT Fatty acids, compounds  
RL: BIOL (Biological study)  
(C10-15, esters, with sugars, sustained-release tablets containing ibuprofen and)

IT Alcohols, biological studies  
Fatty acids, biological studies  
RL: BIOL (Biological study)  
(C10-25, sustained-release tablets containing ibuprofen and sucrose fatty ester and)

IT Carbohydrates and Sugars, esters  
RL: BIOL (Biological study)  
(esters, with fatty acids, sustained-release tablets containing ibuprofen and)

IT 57-50-1D, Sucrose, monoesters with fatty acids 26446-38-8, Sucrose monopalmitate  
RL: BIOL (Biological study)  
(sustained-release tablets containing ibuprofen and)

IT 57-11-4, Stearic acid, biological studies 9003-39-8, Poly(vinyl pyrrolidone)  
RL: BIOL (Biological study)  
(sustained-release tablets containing ibuprofen and sucrose fatty ester and)

IT 15687-27-1, Ibuprofen  
RL: BIOL (Biological study)  
(sustained-release tablets containing sucrose fatty ester and)

L1 ANSWER 18 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1986:632341 CAPLUS

DOCUMENT NUMBER: 105:232341

TITLE: The compressional properties of dextrose monohydrate and anhydrous dextrose of varying water contents

AUTHOR(S): Armstrong, N. Anthony; Patel, Anil; Jones, Trevor M.

CORPORATE SOURCE: Welsh Sch. Pharm., UWIST, Cardiff, UK

SOURCE: Drug Development and Industrial Pharmacy (1986),

12(11-13), 1885-901

CODEN: DDIPD8; ISSN: 0363-9045

DOCUMENT TYPE: Journal

LANGUAGE: English

TI The compressional properties of dextrose monohydrate and anhydrous dextrose of varying water contents

AB The effect of moisture on the compressional properties of anhydrous dextrose [50-99-7] and dextrose monohydrate (I) [14431-43-7] was examined. Relations between moisture content and both tablet tensile strength and tablet toughness were evaluated. An increase in the moisture content of anhydrous dextrose produced a corresponding increase in both strength parameters up to the 8.9% moisture level, possibly due to a recrystg. effect. However any further increase in moisture content beyond this point produced a marked reduction in both tablet tensile strength and tablet toughness. For I, any increase in moisture content obtained by exposure to elevated humidities led to a reduction in both tensile strength and toughness. The consolidation of both anhydrous dextrose and I was improved with increasing moisture content, presumably due to a lubrication effect.

ST compression dextrose hydrate; water compression dextrose; tablet

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property compression dextrose  
IT Tablets  
(properties of, moisture content of dextrose effect on)

L1 ANSWER 19 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1973:445761 CAPLUS  
DOCUMENT NUMBER: 79:45761  
TITLE: Comparative evaluation of excipients for direct compression. I  
AUTHOR(S): Bolhuis, G. K.; Lerk, C. F.  
CORPORATE SOURCE: Lab. Pharm. Technol., State Univ., Groningen, Neth.  
SOURCE: Pharmaceutisch Weekblad (1973), 108(22), 469-81  
CODEN: PHWEAW; ISSN: 0031-6911  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB Micrpocryst.  $\alpha$ -cellulose, granular cellulose, microfine cellulose, directly compressible starch, amylose, Ca<sub>3</sub>(PO<sub>4</sub>)<sub>2</sub>.2H<sub>2</sub>O, dextrose monohydrate, spray-crystallized dextrose, anhydrous lactose, and spray dried lactose were evaluated for tabletting by direct compression. Characteristics for direct compression at different pressures were the coefficient of variation of upper punch force, the ratio of lower to upper punch force and the ejection force during compression and ejection, % of total energy input immediately recovered as elastic energy, and the weight variation, crushing strength and disintegration time of the compacts formed.  
ST excipient tablet direct compression  
IT Tablets  
(compression of, excipients in relation to)  
IT 50-99-7, biological studies 63-42-3 7758-87-4 9004-34-6, biological studies 9005-25-8, biological studies 9005-82-7  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(pharmaceutical excipient, compression of, tablet properties in relation to)

L1 ANSWER 20 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1970:491229 CAPLUS  
DOCUMENT NUMBER: 73:91229  
TITLE: Surface area measurements in compressed powder system  
AUTHOR(S): Armstrong, Norman Anthony; Griffiths, Ryland V.  
CORPORATE SOURCE: Inst. Sci. Technol., Univ. Wales, Cardiff, UK  
SOURCE: Pharmaceutica Acta Helvetiae (1970), 45(9), 583-8  
CODEN: PAHEAA; ISSN: 0031-6865  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB The surface areas of dry and moist compacts of phenacetin (I), paracetamol (II) and dextrose monohydrate (III), prepared by compression in a hydraulic press, were determined by N gas adsorption in a continuous-flow system of N and He. As compression pressure is increased in forming the compacts, the surface area rises to a maximum, falls due to bonding between adjacent particles, and then rises again for I and III. Water in the compacts (2.5-6.6%) reduced surface area due to improved lubrication and, for the more soluble II and III, to recrystn. permitting formation of interparticulate bonds.  
IT Tablets  
(surface area of compressed powder systems for)

L1 ANSWER 21 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

09752899

ACCESSION NUMBER: 1969:95610 CAPLUS  
DOCUMENT NUMBER: 70:95610  
TITLE: Chewing-gum products  
INVENTOR(S): Bucher, Robert C.  
PATENT ASSIGNEE(S): Fleer, Frank H., Corp.  
SOURCE: Ger., 8 pp.  
CODEN: GWXXAW  
DOCUMENT TYPE: Patent  
LANGUAGE: German  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 1288246		19690130	DE 1964-F44582	19641201
PRIORITY APPLN. INFO.: US 19631202				
AB Dried, finely divided sugar is added to a molten, essentially water-free chewing-gum base, which has been preheated to 77-121°, and mixed till a dry, crumbling, powdery mixture is obtained. The chewing-gum base is used in a proportion of 5-40 weight %. Thus, in a kettle previously heated at 66-82°, the chewing-gum base (12.5%) at 99° is introduced and mixed with aroma substances, coloring material, and 20% of the sugar (in total, 18.75% dextrose monohydrate (99%), particle size 0.6175 mm., is added). After once more adding 20% of the sugar, and mixing, the rest of the sugar is added and mixed. The warm mixture (54-71°) is agitated in a trough. The product floats in water and is not hygroscopic. The product pieces suitable for chewing-gum are coated with sugar and used as chewing-gum; the rest is pulverized to a particle size of 0.833 mm. and pressed in tablets.				

L1 ANSWER 22 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1967:118865 CAPLUS  
DOCUMENT NUMBER: 66:118865  
TITLE: Prolonged acting pharmaceutical compositions  
INVENTOR(S): Stephenson, Douglas  
PATENT ASSIGNEE(S): Wellcome Foundation Ltd.  
SOURCE: Brit., 5 pp. Addn. to Brit. 906422  
CODEN: BRXXAA  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
GB 1063872		19670330	GB 1962-30868	19631111
AB Addition to Brit. 906,422 (CA 58, 2328g). Tablets for prolonged effect contain a water-soluble drug, a slowly digestible substance, and a hydrophobic waxy binding agent. The core contains procyclidine-HCl (Kemadrin) (I) 2.5, polyethylene glycol 4000 27, and Mg stearate 0.4 mg., to which a middle layer is applied containing I 5.5, hydrogenated castor oil 64, casein 50, and Mg stearate 37 mg., plus an outer layer containing I 2, lactose 118, dextrose monohydrate 70, starch 24.4, and Mg stearate 2.5 mg.				
IT Tablets (sustained-release)				

L1 ANSWER 23 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 1965:438398 CAPLUS  
 DOCUMENT NUMBER: 63:38398  
 ORIGINAL REFERENCE NO.: 63:6800a-c  
 TITLE: Anthelmintic tablets  
 INVENTOR(S): Stephenson, Douglas  
 PATENT ASSIGNEE(S): Wellcome Foundation Ltd.  
 SOURCE: 6 pp.  
 DOCUMENT TYPE: Patent  
 LANGUAGE: Unavailable  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	-----	-----	-----	-----	-----
	GB 994742		19650610	GB 1960-31223	19600909
TI	Anthelmintic tablets				
AB	The preparation of tablets containing anthelmintics of the bephenium type, I, as an inner core and piperazine (II) in the outer coating is described. The coating of II may be uniform in thickness, or thicker on one side than on the other, or carry a depression on one face. The method of manufacture is described. A typical tablet contains as inner portion I (R = H, R' = 2-thienyl) p-chlorobenzenesulfonate 216.25, alginic acid 2.165, potato starch 43.25, and Mg stearate 3.25 mg. The coating contains II phosphate 260, lactose 78, dextrose monohydrate or sucrose 78, potato starch 26, and Mg stearate 5.2 mg. The completed tablet of thickness 5.75 mm. and diameter 12.6 mm. contains a hole in one face of diameter 4-6 mm. and depth 1.5-2 mm. The tablets allow controlled release of the anthelmintic components.				
IT	Anthelmintics (tablets containing)				
IT	Ammonium, dimethyl(2-phenoxyethyl)-2-thenyl, p-chlorobenzenesulfonate (anthelmintic tablet containing)				
IT	14538-56-8, Piperazine, phosphate (anthelmintic tablet containing)				

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(FILE 'HOME' ENTERED AT 15:48:52 ON 04 OCT 2006)

FILE 'CAPLUS' ENTERED AT 15:49:23 ON 04 OCT 2006

L1	23 SEA ABB=ON PLU=ON TABLET AND DEXTROSE MONOHYDRATE
L2	7 SEA ABB=ON PLU=ON TABLET AND DEXTROSE MONOHYDRATE AND (SOFT OR CHEWABLE)
L3	1 SEA ABB=ON PLU=ON L2 AND FAT
L4	1 SEA ABB=ON PLU=ON L2 AND (OIL OR FAT OR LIPID OR FATTY ADJ ACID)
L5	4 SEA ABB=ON PLU=ON L1 AND (OIL OR FAT OR LIPID OR FATTY ADJ ACID)
L6	19 SEA ABB=ON PLU=ON L1 NOT L5
L7	1 SEA ABB=ON PLU=ON (L1 OR L6) AND FAT FREE D L5 IBIB KWIC D L5 IBIB KWIC 1-Y

FILE 'CAPLUS' ENTERED AT 16:15:13 ON 04 OCT 2006

09752899

D L1 IBIB KWIC  
D L1 IBIB KWIC 1-

FILE HOME

FILE CAPLUS

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FILE LAST UPDATED: 3 Oct 2006 (20061003/ED)

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YOU HAVE REQUESTED DATA FROM 19 ANSWERS - CONTINUE? Y/(N):y

L6 ANSWER 1 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 2005:1885 CAPLUS  
DOCUMENT NUMBER: 142:79974  
TITLE: Soft tablet containing high molecular weight cellulosics  
INVENTOR(S): Wynn, David; Parikh, Nick  
PATENT ASSIGNEE(S): USA  
SOURCE: U.S. Pat. Appl. Publ., 8 pp.  
CODEN: USXXCO  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 2  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2004265373	A1	20041230	US 2003-608681	20030627
CA 2472432	AA	20041227	CA 2004-2472432	20040625
EP 1498114	A1	20050119	EP 2004-253844	20040625
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR				
PRIORITY APPLN. INFO.:			US 2003-607766	A 20030627
			US 2003-608681	A 20030627

L6 ANSWER 2 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 2005:993 CAPLUS  
DOCUMENT NUMBER: 142:79963  
TITLE: Soft tablets containing high molecular

lc

INVENTOR(S): weight celluloses  
 Wynn, David; Parikh, Nick  
 PATENT ASSIGNEE(S): USA  
 SOURCE: U.S. Pat. Appl. Publ., 9 pp.  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 2  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2004265372	A1	20041230	US 2003-607766	20030627
CA 2472432	AA	20041227	CA 2004-2472432	20040625
EP 1491184	A1	20041229	EP 2004-253843	20040625
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR				
PRIORITY APPLN. INFO.:			US 2003-607766	A 20030627
			US 2003-608681	A 20030627

L6 . ANSWER 3 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 2004:368929 CAPLUS  
 DOCUMENT NUMBER: 140:363062  
 TITLE: Pharmaceutical compositions of ganciclovir  
 INVENTOR(S): Mathur, Rajeev Shankar; Kumar, Pananchukunnath Manoj;  
 Roy, Sunilendu Bhushan; Malik, Rajiv  
 PATENT ASSIGNEE(S): Ranbaxy Laboratories Limited, India  
 SOURCE: PCT Int. Appl., 21 pp.  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004037263	A1	20040506	WO 2003-IB4664	20031022
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
AU 2003274410	A1	20040513	AU 2003-274410	20031022
EP 1556050	A1	20050727	EP 2003-758391	20031022
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK				
US 2006189565	A1	20060824	US 2006-532024	20060407
PRIORITY APPLN. INFO.:			IN 2002-DE1058	A 20021022
			WO 2003-IB4664	W 20031022
REFERENCE COUNT:	4	THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT		

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L6 ANSWER 4 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 2004:145843 CAPLUS  
DOCUMENT NUMBER: 141:355096  
TITLE: Powdered and granular materials used in the fabrication of compressed tablets  
AUTHOR(S): Delattre, Luc  
CORPORATE SOURCE: Laboratoire de Technologie Pharmaceutique, Departement de Pharmacie, Faculte de Medecine, Universite de Liege, Liege, Belg.  
SOURCE: Bulletin de la Societe Royale des Sciences de Liege (2003), 72(5), 317-339  
CODEN: BSRSA6; ISSN: 0037-9565  
PUBLISHER: Societe Royale des Sciences de Liege  
DOCUMENT TYPE: Journal  
LANGUAGE: French  
REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 5 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 2003:737151 CAPLUS  
DOCUMENT NUMBER: 139:250306  
TITLE: Soft tablets containing high molecular weight polyethylene oxide  
INVENTOR(S): Luber, Joseph; Bunick, Frank J.  
PATENT ASSIGNEE(S): McNeil-PPC, Inc., USA  
SOURCE: U.S. Pat. Appl. Publ., 7 pp.  
CODEN: USXXCO  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2003175336	A1	20030918	US 2002-97000	20020313
US 6753009	B2	20040622		
CA 2421685	AA	20030913	CA 2003-2421685	20030312
PRIORITY APPLN. INFO.:			US 2002-97000	A 20020313
REFERENCE COUNT:	24			

THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 6 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 2003:8105 CAPLUS  
DOCUMENT NUMBER: 138:61356  
TITLE: Method to aid smoking cessation using dextrose and/or levulose  
INVENTOR(S): West, Robert; Hajek, Peter  
PATENT ASSIGNEE(S): UK  
SOURCE: Brit. UK Pat. Appl., 7 pp.  
CODEN: BAXXDU  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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GB 2376885 A1 20021231 GB 2001-15568 20010626  
PRIORITY APPLN. INFO.: GB 2001-15568 20010626

L6 ANSWER 7 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 2002:674573 CAPLUS  
DOCUMENT NUMBER: 137:206554  
TITLE: Chewable tablets containing hydrate excipients.  
INVENTOR(S): Bunick, Frank J.; Luber, Joseph  
PATENT ASSIGNEE(S): McNeil-PPC, Inc., USA  
SOURCE: U.S. Pat. Appl. Publ., 5 pp.  
CODEN: USXXCO  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2002122822	A1	20020905	US 2000-752601	20001229
US 6814978	B2	20041109		
US 2003175339	A1	20030918	US 2003-413804	20030415
			US 2000-752601	A1 20001229

PRIORITY APPLN. INFO.:  
REFERENCE COUNT: 18 THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 8 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 2001:136959 CAPLUS  
DOCUMENT NUMBER: 134:183494  
TITLE: Orally dissolvable prenatal multi-vitamin  
INVENTOR(S): Devries, Tina; Valentine, William; Valentine, William K.  
PATENT ASSIGNEE(S): Warner Chilcott Laboratories Ireland Limited, USA  
SOURCE: PCT Int. Appl., 38 pp.  
CODEN: PIXXD2  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 2  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001011991	A1	20010222	WO 2000-US40557	20000803
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
US 6495177	B1	20021217	US 2000-539850	20000331
PRIORITY APPLN. INFO.:			US 1999-148803P	P 19990813
			US 1999-148806P	P 19990813
			US 2000-539850	A 20000331

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 9 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 1997:471006 CAPLUS  
 DOCUMENT NUMBER: 127:152888  
 TITLE: Potassium carbonate as a desiccant in effervescent tablets  
 AUTHOR(S): Wells, Mickey L.; Wood, Daniel L.; Sanftleben, Ronald;  
 Shaw, Kelley; Hottovy, Jeff; Weber, Thomas; Geoffroy,  
 Jean-Marie; Alkire, Todd G.; Emptage, Michael R.;  
 Sarabia, Rafael  
 CORPORATE SOURCE: Glaxo Wellcome Inc., Research Triangle Park, NC,  
 27709, USA  
 SOURCE: International Journal of Pharmaceutics (1997), 152(2),  
 227-235  
 CODEN: IJPHDE; ISSN: 0378-5173  
 PUBLISHER: Elsevier  
 DOCUMENT TYPE: Journal  
 LANGUAGE: English

L6 ANSWER 10 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 1997:204347 CAPLUS  
 DOCUMENT NUMBER: 126:255506  
 TITLE: Compressed tablet transitory lubricant system  
 INVENTOR(S): Valentine, William; Valentine, William K.  
 PATENT ASSIGNEE(S): Advanced Technology Pharmaceuticals Corporation, USA  
 SOURCE: U.S., 6 pp.  
 CODEN: USXXAM  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5609883	A	19970311	US 1994-307922	19940916
PRIORITY APPLN. INFO.: US 1994-307922				19940916

L6 ANSWER 11 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 1996:155953 CAPLUS  
 DOCUMENT NUMBER: 124:270037  
 TITLE: Using starch in tabletting  
 AUTHOR(S): Vanhemelrijk, J.; Heume, M.  
 CORPORATE SOURCE: Cerestar Euro Centre Food, Vilvoorde, 1800, Belg.  
 SOURCE: Agro-Food-Industry Hi-Tech (1995), 6(5), 9-10  
 CODEN: AIHTEI; ISSN: 1120-6012  
 PUBLISHER: TeknoScienze srl  
 DOCUMENT TYPE: Journal; General Review  
 LANGUAGE: English

L6 ANSWER 12 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 1995:694438 CAPLUS  
 DOCUMENT NUMBER: 123:93108  
 TITLE: Effect of different excipients on release characteristics of acetylsalicylic acid from compressed pellets

09752899

AUTHOR(S): Torrado-Santiago; Torrado, J. J.; Cadorniga, R.  
CORPORATE SOURCE: Fac. Pharm., Complutense Univ., Madrid, Spain  
SOURCE: Pharmazie (1995), 50(7), 476-8  
CODEN: PHARAT; ISSN: 0031-7144  
PUBLISHER: Govi-Verlag Pharmazeutischer Verlag  
DOCUMENT TYPE: Journal  
LANGUAGE: English

L6 ANSWER 13 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1993:678520 CAPLUS  
DOCUMENT NUMBER: 119:278520  
TITLE: "In vitro" drug release of AAS matrix tablets  
AUTHOR(S): Torrado, S.; Torrado, Susana; Torrado V., J.;  
Cadorniga, R.  
COPORATE SOURCE: Univ. Complutense, Madrid, 28040, Spain  
SOURCE: Proc. Int. Symp. Controlled Release Bioact. Mater.,  
20th (1993), 370-1. Editor(s): Roseman, Theodore J.;  
Peppas, Nicholas A.; Gabelnick, Henry L. Controlled  
Release Soc.: Deerfield, Ill.  
CODEN: 59LOAL  
DOCUMENT TYPE: Conference  
LANGUAGE: English

L6 ANSWER 14 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1988:479708 CAPLUS  
DOCUMENT NUMBER: 109:79708  
TITLE: Sustained-release pharmaceutical containing fatty acid  
sugar esters as excipients  
INVENTOR(S): Jansen, Frans Herwigjan; Hendrickx, Jean  
PATENT ASSIGNEE(S): Sanico, N. V., Belg.; N. V. Gantax S. A.  
SOURCE: Eur. Pat. Appl., 13 pp.  
CODEN: EPXXDW  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 230332	A1	19870729	EP 1987-200031	19870112
R: AT, BE, CH, NL 8600050	DE, ES, FR, GB, GR, IT, LI, LU, NL, SE			
FI 8700092	A	19870803	NL 1986-50	19860113
NO 8700104	A	19870714	FI 1987-92	19870112
DK 8700161	A	19870714	NO 1987-104	19870112
ZA 8700216	A	19870826	DK 1987-161	19870113
JP 62209025	A2	19870914	ZA 1987-216	19870113
PRIORITY APPLN. INFO.:			JP 1987-7403	19870113
			NL 1986-50	A 19860113

L6 ANSWER 15 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1986:632341 CAPLUS  
DOCUMENT NUMBER: 105:232341  
TITLE: The compressional properties of dextrose  
monohydrate and anhydrous dextrose of varying  
water contents  
AUTHOR(S): Armstrong, N. Anthony; Patel, Anil; Jones, Trevor M.  
CORPORATE SOURCE: Welsh Sch. Pharm., UWIST, Cardiff, UK

09752899

SOURCE: Drug Development and Industrial Pharmacy (1986),  
12(11-13), 1885-901  
CODEN: DDIPD8; ISSN: 0363-9045  
DOCUMENT TYPE: Journal  
LANGUAGE: English

L6 ANSWER 16 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1973:445761 CAPLUS  
DOCUMENT NUMBER: 79:45761  
TITLE: Comparative evaluation of excipients for direct compression. I  
AUTHOR(S): Bolhuis, G. K.; Lerk, C. F.  
CORPORATE SOURCE: Lab. Pharm. Technol., State Univ., Groningen, Neth.  
SOURCE: Pharmaceutisch Weekblad (1973), 108(22), 469-81  
CODEN: PHWEAW; ISSN: 0031-6911  
DOCUMENT TYPE: Journal  
LANGUAGE: English

L6 ANSWER 17 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1970:491229 CAPLUS  
DOCUMENT NUMBER: 73:91229  
TITLE: Surface area measurements in compressed powder system  
AUTHOR(S): Armstrong, Norman Anthony; Griffiths, Ryland V.  
CORPORATE SOURCE: Inst. Sci. Technol., Univ. Wales, Cardiff, UK  
SOURCE: Pharmaceutica Acta Helvetiae (1970), 45(9), 583-8  
CODEN: PAHEAA; ISSN: 0031-6865  
DOCUMENT TYPE: Journal  
LANGUAGE: English

L6 ANSWER 18 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1969:95610 CAPLUS  
DOCUMENT NUMBER: 70:95610  
TITLE: Chewing-gum products  
INVENTOR(S): Bucher, Robert C.  
PATENT ASSIGNEE(S): Fleer, Frank H., Corp.  
SOURCE: Ger., 8 pp.  
CODEN: GWXXAW  
DOCUMENT TYPE: Patent  
LANGUAGE: German  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 1288246	-----	19690130	DE 1964-F44582	19641201
PRIORITY APPLN. INFO.:			US	19631202

L6 ANSWER 19 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN  
ACCESSION NUMBER: 1965:438398 CAPLUS  
DOCUMENT NUMBER: 63:38398  
ORIGINAL REFERENCE NO.: 63:6800a-c  
TITLE: Anthelmintic tablets  
INVENTOR(S): Stephenson, Douglas  
PATENT ASSIGNEE(S): Wellcome Foundation Ltd.  
SOURCE: 6 pp.  
DOCUMENT TYPE: Patent  
LANGUAGE: Unavailable

09752899

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
-----	-----	-----	-----	-----
GB 994742		19650610	GB 1960-31223	19600909

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(FILE 'HOME' ENTERED AT 15:48:52 ON 04 OCT 2006)

FILE 'CAPLUS' ENTERED AT 15:49:23 ON 04 OCT 2006

L1	23	SEA ABB=ON	PLU=ON	TABLET AND DEXTROSE MONOHYDRATE
L2	7	SEA ABB=ON	PLU=ON	TABLET AND DEXTROSE MONOHYDRATE AND (SOFT OR CHEWABLE)
L3	1	SEA ABB=ON	PLU=ON	L2 AND FAT
L4	1	SEA ABB=ON	PLU=ON	L2 AND (OIL OR FAT OR LIPID OR FATTY ADJ ACID)
L5	4	SEA ABB=ON	PLU=ON	L1 AND (OIL OR FAT OR LIPID OR FATTY ADJ ACID)
L6	19	SEA ABB=ON	PLU=ON	L1 NOT L5
L7	1	SEA ABB=ON	PLU=ON	(L1 OR L6) AND FAT FREE D L5 IBIB KWIC D L5 IBIB KWIC 1-Y